```
Description
Set
       Items
               (HLA OR (HUMAN(W)LEUKOCYTE(W)ANTIGEN?)) (2N) (G (W) (2 OR 3
S1
          16
             OR 4 OR 5 OR 6))
S2
           7
               RD (unique items)
               ((HLA OR (HUMAN(W)LEUKOCYTE(W)ANTIGEN?)) (2N) (G)) (5N) (I-
S3
         150
            SOFORM? OR (SPLIC? (W) (ALTERNATIVE? OR VARIANT?)))
               S3 NOT PY>1997
S4
          39
               RD (unique items)
S5
          18
S6
        1513
               MONOPHOSPHORYL (W) LIPID (W) A
         825
               S6 NOT PY>1995
S7
S8
               S7 (S) (VECTOR? OR PLASMID? OR DNA)
          3
               RD (unique items)
S9
           1
         308
               S7 (S) (ADJUVAN?)
S10
          78
               S10 AND (GENE? OR DNA OR CONSTRUCT?)
S11
S12
          34
               RD (unique items)
S13
          34
               S12 NOT S9
S14
        1587
               ((NAKED(W)DNA) OR PLASMID? OR VECTOR?) (4N) (LIPID?)
S15
         358
               S14 NOT PY>1995
S16
          26
               S15 (S) (GOLD OR PARTICLE?)
S17
          8
               RD (unique items)
S18
         332
               S15 NOT S16
S19
           0
               S18 AND GOLD
S20
         108
               GOLD (S) LIPID? (S) (DNA OR VECTOR? OR PLASMID? OR CONSTRU-
            CT?)
S21
          19
               S20 NOT PY>1995
S22
          7
               RD (unique items)
S23
          37
               GOLD (S) LIPOSOME? (S) (DNA OR VECTOR? OR PLASMID?)
S24
          13
               RD (unique items)
S25
           2
               S24 NOT PY>1995
S26
           0
               LIPID? (4N
S27
       11863
               (DNA OR VECTOR? OR PLASMID?) (4N) LIPID?
S28
        5418
               S27 NOT PY>1995
S29
        109
               S28 (S) (VACCIN? OR IMMUN?)
               RD (unique items)
S30
          65
               S30 AND PLASMID?
S31
          12
               S30 NOT S31
S32
         53
               S32 AND VECTOR?
S33
          3
S34
         53
               S32 NOT S3
S35
          50
               S32 NOT S33
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A novel series of amphiphilic imidazolinium compounds for in vitro and in vivo gene delivery.

Solodin I; Brown CS; Bruno MS; Chow CY; Jang EH; Debs RJ; Heath TD School of Pharmacy, University of Wisconsin, Madison 53706, USA.

Biochemistry (UNITED STATES) Oct 17 1995, 34 (41) p13537-44, ISSN 0006-2960 Journal Code: AOG

Contract/Grant No.: CA58914, CA, NCI; DK45917, DK, NIDDK; HL53762, HL, NHLBI; +

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Document type: Journal Article

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We have developed three catioinic amphiphiles based on the structure 1-[2-(acyloxy)ethyl]-2-alkyl(alkenyl)-3-(2-hydroxyethyl)imidazolinium chlor ide. Although these three compounds differ only in the structure of the hydrophobic acyl chains, they differ greatly in their ability to mediate in vivo and in vitro gene delivery. Moreover, in vitro efficiency is not predictive of in vivo efficiency. The myristoyl form is the most effective compound in vitro, and the oleoyl form is the most effective compound in The compounds readily form suspensions in aqueous media, both in the pure form and as mixtures with either cholesterol dioleoylphosphatidylethanolamine. These suspensions can be sonicated to produce smaller particles . Particle size, electron microscopy, and the ability to capture glucose suggest that these lipids form liposomes on suspension in aqueous media. When mixed with plasmid DNA, the lipid particles appear to fuse and form larger particles . Fusion is maximal critical DNA:lipid ratio where extensive aggregation and precipitation are observed. Therefore, these compounds behave similarly to other cationic liposome-forming lipids upon interaction with DNA.

Cancer gene therapy using plasmid DNA: safety evaluation in rodents and non-human primates.

Parker SE; Vahlsing HL; Serfilippi LM; Franklin CL; Doh SG; Gromkowski SH; Lew D; Manthorpe M; Norman J

Vical Inc., San Diego, CA 92121, USA.

Human gene therapy (UNITED STATES) May 1995, 6 (5) p575-90, ISSN 1043-0342 Journal Code: A12

Comment in Hum Gene Ther. 1995 May;6(5) 549-50; Comment in Hum Gene Ther. 1995 May;6(5):551-2

Languages: ENGLISH

Document type: Journal Article

Record type: Completed

To evaluate the safety of a plasmid DNA -lipid complex, a series of good laboratory practice (GLP) safety studies were conducted with VCL-1005, DNA expression vector containing both the human class I MHC HLA-B7 heavy-chain and the beta 2-microglobulin (beta 2m) light-chain genes formulated with the cationic lipid, DMRIE/DOPE. In mice, the repeated intravenous injection of VCL-1005 at plasmid DNA doses of 0.1, 1.0, or 10 micrograms for 14 days had only incidental effects on clinical chemistry and hematology, and did not result in any organ pathology. Repeated intrahepatic injections of VCL-1005 in mice did not result in significant liver histopathology or significant alterations in liver enzymes. In cynomolgus monkeys, the repeated intravenous administration of VCL-1005 at a cumulative dose of 720 micrograms of DNA had no effects on clinical chemistry, hematology, or organ pathology. Thus, systemic administration of a **plasmid** DNA expression vector containing the coding sequence for a foreign MHC class I molecule did not result in significant toxicity or a pathological immune response in animals. These results suggest that the direct transfer of VCL-1005, a plasmid DNA -lipid complex, could be used for the safe in vivo delivery of recombinant DNA for a cancer gene therapy trial.

Combined experience from Phase I studies with Allovectin-7, a direct gene transfer immunotherapeutic, in patients with metastatic solid tumors (Meeting abstract).

Nabel GJ; Chang AE; Hersh EM; Vogeizang NJ; Rubin J; Silver H; Stah S; Schreiber AB

Univ. of Michigan Med Center, Ann Arbor, MI

Proc Annu Meet Am Soc Clin Oncol; 14 1995 ISSN 0732-183X

Languages: ENGLISH

Document Type: MEETING ABSTRACTS; CLINICAL TRIAL, PHASE I; CLINICAL TRIAL Record type: Completed

Gene therapy provides a novel approach to increase tumor cell recognition by the immune system through the introduction of immunogenic gene products. We report here the combined experience from Phase I trials intralesional injections of Allovectin-7 in patients with metastatic melanoma, renal cell or colorectal carcinoma. The gene transfer product is a highly purified DNA plasmid driving the expression of encoded sequences for the major histocompatibility complex MHC HLA B-7 heavy chain and beta2 macroglobulin formulated with cytofectin (cationic vector , DMRIE/DOPE) . Expression of the class I MHC protein on the tumor cell surface is intended to enhance recognition of putative tumor associated antigens and elicit a systemic cell mediated immune response which may induce tumor regression. Eligibility criteria include adequate performance status (greater than 70%) and organ function, at least 2 measurable lesions and normal lymphocyte response to PHA stimulation. As of November 1994, 39 patients with either metastatic melanoma (18), renal cell or colon carcinoma (11) were treated at 5 clinical centers out of a total of 72 patients to be enrolled. Dosing regimens comprise escalating single doses or multiple injections of 3 ug to 250 ug Allovectin-7. Injected sites have included subcutaneous nodules, regional lymph nodes and under ultrasound or radiographic guidance lung, liver, mediastinal, renal, periaortic, retrocaval and pancreatic masses. No clinical or laboratory toxicities attributable to the gene transfer product have been observed. Adverse events occurred in 8 out of 39 patients and were attributed either to the underlying disease (2 out of 8) or to the injection and study required biopsy procedures (2 pain, 2 hematoma, and 2 small pneumothoraces after injection of lung nodules). Efficiency of gene transfer is followed

by molecular, flow cytometric and immunohistochemistry analyses of biopsies. Immunological responses are followed by serology and cytotoxicity assays and immunocytochemistry of T lymphocytes subset infiltration in tumor biopsies. As of November 1994, partial tumor regression has occurred in 5 of 11 evaluable patients with metastatic melanoma. These studies indicate that intralesional injection of a plasmid DNA /lipid complex can be performed safely and induce tumor regression in some patients.

05991562 EMBASE No: 1995020178

Transfecting neurons and glia in the rat using pH-sensitive immunoliposomes

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Neuroscience Letters ( NEUROSCI. LETT. ) (Ireland) 1995, 184/1 (40-43)

CODEN: NELED ISSN: 0304-3940 DOCUMENT TYPE: Journal; Article

LANGUAGE: ENGLISH SUMMARY LANGUAGE: ENGLISH

Immunoliposomes were constructed using antibody 5-113 (directed to an antigen on the external surface rat glial cells), the antibody Thy 1.1, and a non-immune antibody. The antibodies were conjugated to N-gluytaryl-phosphatidylethanolamine. Liposomes were constructed with these conjugated antibodies, other lipids and a beta-galactosidase plasmid under the control of the cytomegalovirus promoter. When immunoliposomes decorated with one of three different antibodies were injected into the brain or spinal cord of adult rats, the X-gal reaction product was observed in neurons, astrocytes and vascular elements. There was an increase in neuronal labeling when animals were injected with Thy 1.1 conjugated liposomes and there was an increase in glial labeling in animals injected with 5-113 liposomes. In spinal cords, the immunoliposomes appear to penetrate a substantial distance, transfecting neurons several centimeters from the site of delivery. These data suggest that immunoliposomes may provide an effective transfection system for gene delivery in the CNS.

## Cationic lipids direct a viral glycoprotein into the class I major histocompatibility complex antigen-presentation pathway.

Walker C; Selby M; Erickson A; Cataldo D; Valensi JP; Van Nest GV Chiron Corporation, Emeryville, CA 94608.

Proceedings of the National Academy of Sciences of the United States of America (UNITED STATES) Sep 1 1992, 89 (17) p7915-8, ISSN 0027-8424 Journal Code: PV3

Languages: ENGLISH

Document type: Journal Article

Record type: Completed

Recombinant glycoprotein B (gB) of herpes simplex virus (HSV) was processed and presented by class I major histocompatibility complex (MHC) molecules after delivery into cells by using N-[1-(2,3-dioleoyloxy)propyl]-N, N, N-trimethylammonium methyl sulfate (DOTAP), a commercially available used for DNA transfection. Cells treated with cationic lipid DOTAP-associated gB were susceptible to lysis by class I MHC-restricted, HSV-specific cytotoxic T lymphocytes (CTL), and the treated cells memory gB-specific CTL activity in spleen cells from restimulated HSV-infected mice. gB-specific CTL responses were detected in mice with recombinant gB and DOTAP but not in those receiving gB immunized emulsified in complete Freund's adjuvant. Thus, cationic lipids may facilitate induction of CD8+ T-cell responses in vaccinations with recombinant antigens, and they may serve as readily available reagents for dissecting class I MHC immunity to viruses and other intracellular pathogens.